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(FILE 'HOME' ENTERED AT 09:43:04 ON 16 FEB 2004)

INDEX 'ADISCTI, ADISINSIGHT, ADISNEWS, AGRICOLA, ANABSTR, AQUASCI, BIOBUSINESS, BIOCOMMERCE, BIOSIS, BIOTECHABS, BIOTECHDS, BIOTECHNO, CABA, CANCERLIT, CAPLUS, CEABA-VTB, CEN, CIN, CONFSCI, CROPB, CROPUS, DISSABS, DDFB, DDFU, DGENE, DRUGB, DRUGMONOG2, ...' ENTERED AT 09:43:33 ON 16 FEB 2004

SEA HUMAN KINASE

1 FILE BIOCOMMERCE
129 FILE BIOSIS
104 FILE BIOTECHABS
104 FILE BIOTECHDS
30 FILE BIOTECHNO
1 FILE CABA
15 FILE CANCERLIT
155 FILE CAPLUS
2 FILE CEABA-VTB
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1 FILE PHARMAML
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36 FILE SCISEARCH
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261 FILE USPATFULL
49 FILE USPAT2
149 FILE WPIDS
149 FILE WPINDEX

L1

QUE HUMAN KINASE

SEA F5-F20

0* FILE DGENE

FILE 'CAPLUS, WPIDS, BIOSIS, BIOTECHDS, USPAT2, MEDLINE, SCISEARCH, EMBASE, BIOTECHNO, ESBIOWBASE, LIFESCI, TOXCENTER, CANCERLIT, PROMT'
ENTERED AT 09:46:28 ON 16 FEB 2004

L2 521 S L1 AND (INHIBIT? OR MODULAT? OR AGONIST OR ANTAGONIST)
L3 152 S L2 AND AGONIST
L4 98 DUP REM L3 (54 DUPLICATES REMOVED)
L5 175 S L2 AND ANTAGONIST
L6 112 DUP REM L5 (63 DUPLICATES REMOVED)

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L4 ANSWER 96 OF 98 CAPLUS COPYRIGHT 2004 ACS on STN
 ACCESSION NUMBER: 1997:640772 CAPLUS
 DOCUMENT NUMBER: 127:304772
 TITLE: A kinase capable of site-specific phosphorylation of I κ B- α and its regulation and use in the diagnosis or treatment of NF- κ B-related diseases
 INVENTOR(S): Chen, Zhijian J.
 PATENT ASSIGNEE(S): Proscript, Inc., USA
 SOURCE: PCT Int. Appl., 141 pp.
 CODEN: PIXXD2
 DOCUMENT TYPE: Patent
 LANGUAGE: English
 FAMILY ACC. NUM. COUNT: 2
 PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
WO 9735014	A1	19970925	WO 1997-US4195	19970319
W: AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, CA, CH, CN, CU, CZ, DE, DK, EE, ES, FI, GB, GE, GH, HU, IL, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MD, MG, MK, MN, MW, MX, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, TJ, TM, TR, TT, UA, UG, UZ, VN, YU, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM				
RW: GH, KE, LS, MW, SD, SZ, UG, AT, BE, CH, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF, CG, CI, CM, GA, GN, ML, MR, NE, SN, TD, TG				
CA 2249551	AA	19970925	CA 1997-2249551	19970319
AU 9725317	A1	19971010	AU 1997-25317	19970319
AU 728679	B2	20010118		
EP 910645	A1	19990428	EP 1997-916786	19970319
R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, FI				
BR 9710408	A	19990817	BR 1997-10408	19970319
JP 2000510328	T2	20000815	JP 1997-533584	19970319
PRIORITY APPLN. INFO.:			US 1996-616499	A 19960319
			WO 1997-US4195	W 19970319

AB A large multi-subunit protein kinase that catalyzes the site-specific phosphorylation of the NF- κ B inhibitor I κ B α after activation by ubiquitylation by E1 and E2 enzymes is described. The enzyme has 10 subunits of 31, 33, 36, 38, 40, 43, 50, 55, 62, 70, and 85 kilodaltons and phosphorylates I κ B α at residues 32 and 36. CDNAs encoding the subunits of the enzyme may be cloned and characterized for use in the manufacture of the enzyme, e.g. for preparation of the enzyme

for

assays or preparation of antibodies for diagnostic or therapeutic use. The cDNAs may be used to develop probes to detect and quantify gene expression or in gene therapy. Agonist or antagonist ligands of the kinase may be of diagnostic or therapeutic use. More specifically, this invention relates to selective inhibitors of the kinase and to structure-based design of ligands, agonists, and antagonists of the kinase.

L4 ANSWER 97 OF 98 CAPLUS COPYRIGHT 2004 ACS on STN

ACCESSION NUMBER: 1993:253031 CAPLUS

DOCUMENT NUMBER: 118:253031

TITLE: Stimulation of MHC class I transcription by interferon- γ involves a non-A, non-C kinase in addition to protein kinase C

AUTHOR(S): Radford, James E., Jr.; Waring, Jeffrey F.; Pohlman, Joyce K.; Ginder, Gordon D.

CORPORATE SOURCE: Inst. Hum. Genet., Univ. Minnesota, Minneapolis, MN, 55455, USA

SOURCE: Journal of Interferon Research (1993), 13(2), 133-41
CODEN: JIREDJ; ISSN: 0197-8357

DOCUMENT TYPE: Journal

LANGUAGE: English

AB The signal pathways by which interferon- γ (IFN- γ) is able to up-regulate major histocompatibility complex (MHC) class I transcription were studied in two human hematopoietic tumor cell lines, K562 and Ramos. These studies suggest that the IFN- γ signal is transduced via an H7- and staurosporine-sensitive kinase that is distinct from protein kinase C (PKC) and protein kinase A (PKA) in both cell types. Ramos cells appear to utilize an addnl. pathway involving double-stranded RNA-dependent protein kinase. PKC and possibly PKA appear to be involved in one or more intersecting pathways by which agonists of these kinases are able to act synergistically with IFN- γ , but activation of these latter pathways is neither necessary nor sufficient for induction of MHC class I transcription. Modulation of G-protein- and Ca²⁺-calmodulin-associated pathways and arachidonic acid metabolism had no effect

on constitutive or IFN- γ -stimulated class I transcription. The class I stimulatory factor produced in response to IFN- γ treatment appears to have a short t_{1/2}. The identity of this factor is unknown, but is likely to be distinct from known mediators of IFN-stimulated transcription. Gene and cell-type specificity in the signal transduction pathways utilized by IFN- γ implies that such pathways may be useful targets for exptl. and therapeutic manipulation.